

Development of an *in Vitro* Screening Test To Evaluate the *In Vivo* Bioaccessibility of Ingested Mine-Waste Lead

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A screening-level *in vitro* test was developed to evaluate the relative solubility of ingested lead (Pb) from different mine wastes in the gastrointestinal (GI) tract. The *in vitro* method, modeled after assay methods for available iron from food, used a laboratory digestion procedure designed to reproduce GI tract chemistry and function. The *in vitro* method was independently calibrated against a rabbit feeding study, demonstrating that only 1-6% of the total Pb in four mine-waste samples with disparate Pb mineralogy was bioaccessible. *In vitro* method development tests indicated that H⁺ concentration and Cl⁻ complexation control dissolution of Pb-bearing minerals in the stomach and that both GI tract enzymes and organic acids are necessary to maintain Pb in the soluble form on entering the small intestine. The experimental results indicate that ingestion of Pb-bearing mine wastes results in limited Pb solubility and that the *in vitro* test provides a screening-level estimate of the maximum available Pb from mine wastes.

Introduction

When assessing risks associated with lead (Pb)-contaminated soils, one exposure pathway typically evaluated is soil ingestion by children. Standard procedures recommended by the U.S. Environmental Protection Agency (EPA) for estimating soil Pb exposures using the Uptake Biokinetic Model assume that a typical child will ingest 100 mg of soil/day and that 30% of the ingested Pb will be absorbed into the systemic circulation (i.e., will be bioavailable) (1). However, recently completed animal studies indicate that lead from mine waste is much less bioavailable than lead from other sources (2-4). Our previous research has demonstrated that Pb-bearing minerals dissolve slowly and incompletely in the gastrointestinal (GI) tract, due to a variety of geochemical factors (5) resulting in Pb forms that are bioaccessible. The limited dissolution of Pb-bearing minerals in the GI tract is most likely responsible for the low bioavailability observed in the animal studies, because Pb absorption has been shown to occur from the soluble phase in the small intestine (6, 7).

The geochemical factors controlling dissolution of metals from mine waste include the mineral composition, the degree of encapsulation, and the presence of alteration rinds. Because the solubility of Pb from mine waste is

Table I. Occurrence of Pb Minerals in Mine-Waste Samples

Pb phase	sample no.			
	BMW-1	BMW-2	BMW-3	BMW-4
anglesite (PbSO ₄) (%)	59	4	10	
galena (PbS) (%)	26		1	5
lead phosphates ^a (%)	4	7	37	39
manganese-lead oxides ^b (%)		31	7	26
iron-lead oxides ^c (%)	2	29	6	
iron-lead sulfates ^d (%)	2	24	39	
lead oxides [PbO and Pb(OH) ₂] (%)	1	3		17
lead metallica ^e (%)				2
lead silicates (PbSiO ₃ and PbSiO ₄) (%)		2		11
lead barite [(Ba-Pb)SO ₄] (%)	3			
cerussite (PbCO ₃) (%)	3			
Pb concn (mg/kg) ^f	3900	1030	5820	1790
As concn (mg/kg) ^f	1380	620	1180	420
Fe concn (%) ^f	6.9	12.3	5.0	5.4
Mn concn (mg/kg) ^f	2200	2000	1100	1200
Zn concn (mg/kg) ^f	6040	NA ⁱ	NA	1600
Ca concn (%) ^f	0.7	2.9	1.2	0.8
pH ^h	3.7	2.8	3.8	2.6

^a Lead phosphate grains contain variable compositions of Pb, PO₄, SO₄, and halogens. ^b Manganese-lead oxide particles contain more Mn than Pb, with variable compositions of constituents. ^c Iron-lead oxide particles contain more Fe than Pb, with variable compositions of constituents. ^d Iron-lead sulfate grains contain more Fe than Pb, with variable compositions of constituents. ^e Lead metallica contain elemental Pb with Fe, Mn, As, Cu, and Zn in varying proportions. ^f Determined by digestion (31) and atomic absorption spectroscopy (31). ^g Determined by X-ray fluorescence. ^h Determined by the saturated paste method (32). ⁱ NA, not analyzed.

constrained by the composition of the mineral assemblage, site-specific mineralogical data are necessary to characterize the availability of Pb (2, 5). Two dominant Pb mineral assemblages—sulfide/sulfate and oxide/phosphate—have been identified in residential soils containing mine waste (8). The sulfide/sulfate assemblage consists of galena (PbS) grains altering to the oxidation product anglesite (PbSO₄) or enclosed in pyrite (FeS₂) or silicate (SiO₂) matrices, making them unavailable for alteration or dissolution. Precipitation of jarosite [KFe₃(SO₄)₂(OH)₆] rinds on Pb-bearing particles was also observed and would retard Pb dissolution, both physically by reduction in the exposed surface area and chemically due to the insoluble nature of jarosite in acidic (pH < 4) media (9). The oxide/phosphate assemblage consists of lead phosphates of varying compositions, lead ferromanganese oxides, iron-lead sulfates that are similar in composition to the mineral lead jarosite [PbFe₆(SO₄)₄(OH)₁₂], and lead oxides [PbO and Pb(OH)₂, Table I]. These Pb minerals occur in mine waste and soils as a complex set of alteration products

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that are weathering toward more stable lead phosphates. The geochemical controls on Pb dissolution from one of the mine-waste samples (BMW-1) used in this study have been discussed in detail in previous papers (5, 10).

The mass of Pb dissolved is also controlled by physiological factors, including gastric pH, the rate of Pb dissolution relative to the residence time of Pb-bearing mine waste in the GI tract, and by *in vivo* sorption and precipitation reactions that may limit dissolved Pb concentrations. Gastric pH, which in humans ranges from a basal level of 1–2 to 4–6 after ingestion of food, with its attendant buffering capacity (11, 12), appears to be one of the most important physiological factors in determining the mass of Pb solubilized from a mine waste. Lead that enters the fluid phase in a fasting stomach (assumed pH = 1.3 and high chloride activity) will be present at approximately equimolar concentrations of Pb^{2+} and $PbCl^+$ (13). Another critical factor is the residence time in the stomach. The gastric contents are emptied completely into the small intestine within approximately 2 h in humans when fed various test meals (14).

On entering the duodenum, $NaHCO_3$ excreted with the pancreatic juices is mixed with the intestinal chyme, resulting in an increase in pH to approximately 7 in humans (15). Consequently, dissolved Pb concentrations are likely to decrease in the small intestine due to adsorption to mineral and food-particle surfaces with increasing pH and as a result of precipitation reactions (16). For the purpose of this discussion, the fluid or soluble phase within the small intestine is operationally defined as the fraction that can be separated from the solid fraction by relatively low-speed centrifugation, as defined in the methods section. Lead in the soluble small intestinal phase will be distributed between fractions bound by proteins and enzymes (17, 18); complexed by amino acids (19, 20), low molecular weight carboxylic acids (21), and tannins and humic acids released from ingested soil and food, respectively (16); adsorbed to suspended particulates in the fluid phase; or present as free lead cations and hydroxide complexes. The distribution of Pb between these forms will control the solubility and, consequently, the mass available for absorption across the intestinal epithelium.

Because of the complexity of the geochemical and physiological factors controlling dissolution of lead from mine waste, it was determined that a rapid, inexpensive *in vitro* method was needed to investigate interactions of the various factors and to compare the relative solubility of Pb from solid sources. The *in vitro* method for estimating Pb availability presented in this paper was modeled after an *in vitro* assay method for available Fe in foodstuffs (22–26). The Fe availability method simulates the stomach and small intestinal phases of digestion using solutions of specific pH that contain digestive enzymes (pepsin in the stomach, pancreatic enzymes and bile acids in the small intestine) mixed with the test substrate to reproduce GI tract function and chemistry.

The pH values of 1.3 and 7.0, selected for the gastric and small intestinal incubations, respectively, were based on measurements in fasting rabbit stomachs and small intestines. The strongly acidic stomach solution selected for the *in vitro* method is representative of a fasting child and maximizes Pb dissolution to provide an upper-bound estimate of available Pb that would apply to ingestion of small particles due to mouthing behavior by children several hours after food ingestion or under fasting con-

ditions. Selection of appropriate concentrations for digestive enzymes is problematic because concentrations are highly variable in the human system. The enzyme concentrations selected for the *in vitro* test were those used in refs 23 and 24 for *in vitro* estimation of Fe from food. The method developed herein included the addition of small quantities of organic acids that were determined to be present in the rabbit GI tract.

Gastric mixing rate (the rate at which ingested material is mixed with fluid in the stomach due to peristalsis) will also affect mineral dissolution kinetics and influence the concentrations of Pb solubilized in the stomach. Therefore, mixing of the *in vitro* flask contents employed a mixing rate designed to replicate *in vivo* mixing, as previously determined in Pb dissolution rate experiments (9). The test material mass (4 g) and fluid volume (40 mL) for the *in vitro* experiments were based on those found in the stomachs and small intestines of 33 New Zealand White rabbits. The rabbits had been dosed with 2.0 g of mine waste/kg of body weight and weighed, on average, 2.1 kg, resulting in 4.2 g of ingested mine waste (16.4 mg of Pb).

To test the validity of the *in vitro* model, an *in vivo* experiment was conducted to assess the dissolution of Pb from mine waste during passage through the GI tract of New Zealand White rabbits. A mine-waste dose of 2.0 g/kg of body weight was selected to represent the worst-case scenario of a child with pica-for-soil (e.g., a child who intentionally ingests soil), who may ingest up to 10 g of soil/day (27, 28). Although differences in digestive anatomy and function between humans and rabbits, such as coprophagy, biliary excretion, and development of Pb absorption mechanisms during growth, must be considered when assessing Pb bioavailability, the primary factors controlling Pb dissolution (i.e., pH, mixing, and transit time) are comparable for humans and rabbits (5, 11, 12, 14, 29). Therefore, the rabbit GI tract provides an appropriate model to assess Pb bioaccessibility from a mine waste on ingestion by a child. Values for several important parameters in the *in vitro* model were therefore based on observed values in rabbits, as well as humans.

Methods

Mine-Waste Collection and Characterization. The mine waste used for both the *in vivo* experiment and *in vitro* method development (BMW-1) was blended using five mine-waste samples (waste rock and mine overburden material) collected from one mining site to achieve a desired Pb content (3900 mg/kg of Pb). Before blending, each of the five mine-waste samples was air-dried and sieved to <250 μm using a mechanical sieve shaker, because predominantly smaller particles adhere to children's hands and may be ingested (30). The remaining samples (BMW-2–4) were composite samples collected from individual mine-waste piles and treated in an identical manner to BMW-1. Bulk Pb concentrations were determined for each sample by digestion in HNO_3/H_2O_2 , followed by atomic absorption spectroscopy (31). Bulk arsenic (As), Fe, manganese (Mn), zinc (Zn), and calcium (Ca) concentrations were determined by X-ray fluorescence. Sample pH was determined by the saturated paste method

(32). Lead minerals in the mine-waste samples were identified by electron microprobe (JEOL 8600) at the Department of Geological Sciences, University of Colorado at Boulder (Table I). A description of microprobe methods, and more detailed descriptions of the Pb mineralogy, may be found in refs 5 and 8.

In Vivo Experiment. Thirty-six female New Zealand White rabbits, weighing an average of 2.1 kg at 12 weeks, were used in the *in vivo* study undertaken at Battelle Columbus Laboratories (Columbus, OH) and performed in compliance with Good Laboratory Practice Regulations (33). Animals were caged in individual stainless steel cages with mesh bottoms. To avoid interactions between the rabbit chow and the mine waste, all of the rabbits were fasted for 16 h prior to dosing and 4 h after. During nonfasting periods, Purina high-fiber rabbit chow was provided *ad libitum*. Deionized (DI) water was available at all times. Twenty-seven rabbits were dosed with 2.0 ± 0.02 g of BMW-1 (in gelatin capsules)/kg of body weight, and three rabbits each were killed at 0.5, 1, 1.5, 3, 6, 8, 16, 24, and 36 h after dosing. One group of three rabbits was given a mass of soluble Pb salts equal to that given to the mine-waste-dosed animals (14.3 mg of lead acetate $[\text{Pb}(\text{CH}_3\text{CO}_2)_2 \cdot 3\text{H}_2\text{O}]$ /kg of body weight) and killed 1 h after dosing. Two groups of three rabbits each served as controls and were killed at 1 and 36 h after dosing.

Whole blood was collected from all rabbits via cardiac puncture after sedation with sodium pentobarbital. Subsequently, a lethal injection of pentobarbital was administered, the stomach and intestines were exposed, and the region below the pyloric valve was clamped to prevent passage of the stomach contents into the duodenum. Measurements of pH were performed using an Orion portable pH meter in conjunction with a Ross semimicro pH electrode. The electrode was inserted through an incision in the stomach wall for measurement of pH in the fundic and pyloric regions of the stomach. These measurements were also taken in the duodenum and ileum by removing the small intestine and gently inserting the probe into either end of the segment. Large intestine measurements were made in the middle of the cecum.

Samples of the contents of the stomach, small intestine, and large intestine were collected from each animal in 50-mL Corning polypropylene centrifuge tubes and centrifuged at approximately 2100g for 25 min in a Beckman Model TJ6 centrifuge. Approximately 30–40 mL of stomach material was collected with a spoon through a slit in the stomach wall, and any remaining material was removed from the stomach with a spoon, collected into a disposable beaker, and weighed. The volume of material in the fasted rabbit stomachs was approximately 30 mL, while the volume in fed rabbit stomachs was approximately 60 mL. Entire small intestinal contents (10–20 mL) were extruded into single centrifuge tubes. Approximately 40 mL of large intestinal material was collected with a spoon through a slit in the middle portion of the organ, and the remaining material was collected with a spoon into a disposable beaker and weighed. The fluid phase was decanted, and reagent-grade HNO_3 (17N) was used to acidify the fluid samples (1% v/v). Solid samples were dried to determine percent moisture and digested in $\text{HNO}_3/\text{H}_2\text{O}_2$ (31). A Thermo Jarrel Ash, Video 11E atomic absorption spectrophotometer, equipped with a Model CTF 188 graphite furnace and Smith-Hieftje background correction, was used for analysis of samples with <2 mg/L

of Pb, and the same instrument, equipped for flame analysis, was used for samples with >2 mg/L of Pb. Organic acid analyses were performed by capillary electrophoresis at Galbraith Labs (Knoxville, TN).

Analytical quality assurance and quality control (QA/QC) samples consisted of 1 in 20 samples analyzed as matrix spikes and duplicates, calibration verification, continuing calibration verification and blanks, and the use of the method of standard additions when recalcitrant matrices were encountered. Based on quality control limits for acceptable analytical results (34), all data were accurate and precise.

In Vitro Experiments. A mine-waste/solution ratio of 1:10 (4.0 ± 0.01 g of mine waste and 40 mL of fluid) was selected for the *in vitro* method because each rabbit was dosed with 2 g of BMW-1/kg of body weight (average of 4.2 g of BMW-1/rabbit), and the 1:10 solid-to-fluid ratio was the average ratio observed in the 36 rabbit stomachs and small intestines. The *in vivo* experiment was designed to minimize interactions between the mine waste and rabbit chow; however, because rabbits practice coprophagy (35), some interaction between mine waste and food is inevitable. Therefore, *in vitro* tests were conducted with both mine waste alone (in triplicate) and a mine-waste/rabbit chow mixture (4.0 g of BMW-1, 1.0 g of rabbit chow, 40 mL of fluid), in duplicate. The *in vitro* test was also conducted with an equivalent mass of Pb as lead acetate, in both the presence and absence of Pb-free mine waste, to allow calculation of Pb solubility from BMW-1 relative to lead acetate. Subsequent to method development using BMW-1, three additional mine-waste samples (BMW-2–4, Table I) were subjected to the *in vitro* test to examine Pb availability from mine waste with disparate Pb mineralogies.

A total of 40 mL of type I DI water was adjusted to pH 1.3 with HCl. Mine waste (4 g), pepsin (50 mg, activity of 800–2500 units/mg, Sigma Chemical Co.), and organic acids (acetate [0.5 mL], citrate [0.5 g], lactate [0.42 mL], and malate [0.5 g]) were added, and the flask was placed on a wrist-action shaker in a water bath at 37 °C, representative of GI tract temperature (23). The pH was checked at 10-min intervals during the first half hour and at half-hour intervals thereafter, and measured volumes of 12N HCl were added to maintain a pH of 1.3. Aliquots (2 mL) were removed from the reaction flask at 0.5, 1.0, 1.5, and 2.0 h after initiation of the reaction and centrifuged immediately at approximately 2100g for 25 min, and the liquid fraction was decanted for Pb analysis. After 2 h, the reaction vessel was titrated to $\text{pH } 7.0 \pm 0.2$ by the addition of a dialysis bag (8000 MWCO, Spectra/Por cellulose ester tubing) containing 3 mL of DI and NaHCO_3 of equivalent molarity to the calculated acidity (HCl) in the flask. The system required approximately 30 min to reach a pH of 7.0 ± 0.2 , after which the dialysis bag was removed and the contents were emptied into the reaction flask. Pancreatin (20 mg, activity equivalent to $4 \times$ U.S. Pharmacopeia specifications, Sigma Chemical Co.) and bile extract (70 mg) were added, and the flask was returned to the water bath. Two hours were allowed for the flask contents to reach equilibrium, the fluid volume was measured, and a 10-mL fluid sample was collected by centrifugation for Pb analysis. Fluids were analyzed for Pb using the methods and instrumentation described in the *in vivo* experimental section. GI tract enzymes and

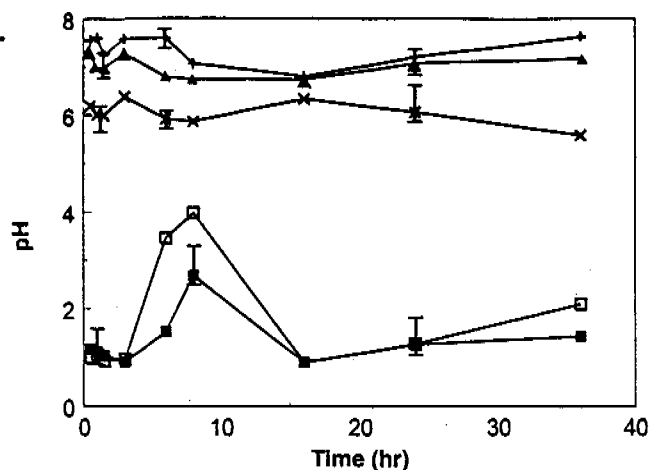


Figure 1. pH variations in rabbit GI tract as a function of time and location: (■) fundic stomach, (□) pyloric stomach, (▲) duodenum, (+) ileum, (X) cecum. Error bars indicate standard deviation for the measured values in three rabbits.

acids were obtained from Sigma Chemical Co. (St. Louis, MO).

Method development testing for the *in vitro* test also included omission of the organic acids, omission of the GI tract enzymes, and omission of both the acids and enzymes (each test in duplicate). A parallel procedure was carried out (in duplicate) with an equimolar amount of a soluble lead acetate [$\text{Pb}(\text{CH}_3\text{CO}_2)_2 \cdot 3\text{H}_2\text{O}$, 24.5 mg] to evaluate Pb availability from a soluble salt with no matrix present. Finally, the *in vitro* test was conducted with lead acetate (in duplicate) in reaction flasks containing 4 g of low-Pb mine waste (100 mg/kg of Pb). This mine waste was also subjected to the *in vitro* procedure (in duplicate) to determine available Pb. An experimental blank (no solid material in flask) was carried through the procedure during each experiment.

Results

Mineralogy, Metal Concentrations, and Sample pH. Lead mineralogy in samples BMW-1-4 (Table I) was calculated as percent occurrence of Pb-bearing grains and indicates a diverse Pb mineralogy consisting of both sulfate/sulfide and oxide/phosphate phases, consistent with Pb mineralogy observed during a comprehensive study at this site (8). Bulk metals data are presented (Table I) for comparison of these mine-waste samples with samples that may be analyzed by the *in vitro* method in the future. Although the bulk concentrations of As, Fe, Mn, Zn, and Ca will have a minimal effect on the dissolution of Pb-bearing phases, the presence of soluble Fe and Ca will decrease the absorption of Pb across the intestinal epithelium (16). The acidic pH values determined in these samples (2.6–3.8) are consistent with the high Fe concentrations (5.0–12.3 wt %) and the presence of pyrite and suggest that these materials will have a limited capacity to buffer stomach acids on ingestion.

In Vivo pH. The pH measurements collected in the rabbit GI tracts (36 rabbits) indicate that a pH in the range of 1.0–1.3 exists in the stomach under fasting conditions, increasing to 2.8 and 4.1, respectively, in the fundic and pyloric regions of the stomach on ingestion of food (Figure 1). Small and large intestinal pH values were independent of nutritional status and averaged 6.9 in the duodenum, 7.4 in the ileum, and 5.9 in the cecum. These

Table II. Concentrations of Organic Acids in Rabbit GI Tract

organic acid	pK_a	lead-acid log K_{ML} ^a ionic strength 1.0	stomach concn ^a (mg/L)	small intestine ^a (mg/L)
acetate	4.7 ^b	2.1	189	212
butyrate	4.9 ^c	2.1	<10	<10
citrate	pK_1 3.1 ^b	4.1	534	<10
	pK_2 4.8 ^b			
lactate	3.8 ^b	2.0	174	972
malate	pK_1 2.8 ^d	2.4	1302	100
	pK_2 5.7 ^d			
succinate	pK_1 4.2 ^d	2.4	64	<10
	pK_2 5.6 ^d			
tartrate	pK_1 3.1 ^b	2.6	<10	<10
	pK_2 4.8 ^b			

^a Determined by capillary electrophoresis. ^b Ref 40. ^c Ref 41. ^d Ref 42. ^e Ref 36.

values are in agreement with the basal gastric pH of 1.0–2.0 (11, 12) and with the small intestinal pH of 4.7–7.2 (15) for human subjects, validating the use of female New Zealand White rabbits as a GI model for humans with respect to pH. Based on observations in New Zealand White rabbits, pH levels of 1.3 and 7.0 were selected to represent the stomach and small intestinal phases, respectively, in the *in vitro* model.

Organic Acids. Organic acids are known to chelate Pb (21) and may increase solubilized Pb concentrations in the small intestine, promoting transport of Pb across the intestinal epithelium (6). Therefore, concentrations of organic acids were measured by capillary electrophoresis (Table II) in the stomach and small intestinal fluids of a rabbit from the 6-h post-dose group. The pK_a values for the first proton released from all of the carboxylic acids for which analyses were performed are ≥ 2.8 (Table II); therefore, these acids will be protonated during fasting stomach conditions (pH 1–2) but may chelate Pb in the stomach in the presence of food (pH 4–6) and in the small intestine (pH 7). Comparison of stability constants (determined at 25 °C and solution ionic strength of 1.0 (36) for the lead carboxylate complexes) indicates that citrate, if present, will form the most stable complex with Pb ($\log K_{ML} = 4.1$), while the remaining acids will form complexes of decreasing stability in the order tartrate > malate, succinate > acetate, butyrate > lactate. The ability of organic acids to serve as Pb chelators depends on the ability of the soluble chelator to complex Pb (which is pH dependent due to proton competition for the ligand), the strength of Pb^{2+} binding to individual ligands (e.g., selectivity for Pb), and the activity of competing cations (e.g., Ca^{2+} , Zn^{2+} , or Cu^{2+}) that may displace Pb. Therefore, in the small intestine at pH 7, the K_{ML} values for organic acids (Table II) indicate that nearly all Pb complexed by organic acids would be present as lead citrate complexes; however, this does not eliminate the possibility that Pb would be preferentially complexed by stronger chelators (e.g., enzymes and amino acids) or displaced by competing cations.

Malate and citrate were the most concentrated carboxylic acids in the rabbit stomachs, whereas lactate and acetate were most concentrated in the small intestine (Table II). The decrease in citrate and malate concentrations between the stomach and small intestinal fluid may be due to the absorption of these species in the small intestine or to the formation of stable complexes that are

Table III. Pb Mass (μg) Recovered from Rabbit Fluids and Solids, Reported as Arithmetic Mean, Standard Deviation of Three Samples^a

time (h)	stomach fluid ^b	small int. fluid ^b	large int. fluid ^b	blood ^d	stomach solids	small int. solids	large int. solids	% total Pb in stomach in fluid phase ^h (%)
0.5	230 \pm 170	6.9 \pm 6.7	3.0 ^c	2.0 \pm 0.4	6980 \pm 2290	38 \pm 20	170 \pm 60	3 \pm 2
1	260 \pm 140	28 \pm 34	3.8 \pm 2.4 ^c	1.5 \pm 0.5	6670 \pm 2820	480 \pm 710	130 \pm 47	4 \pm 2
1.5	620 \pm 250	140 \pm 27 ^c	—	1.7 \pm 0.4	6210 \pm 2070	1230 \pm 630	690 \pm 760	9 \pm 4
3	62 \pm 53	250 \pm 220	32 \pm 25	5.4 \pm 2.8	850 \pm 30	4580 \pm 4830	550 \pm 530	7 \pm 6
6	5.8 \pm 3.8	6.1 \pm 6.7	53 \pm 18	7.1 \pm 2.1	62 \pm 80	160 \pm 230	1170 \pm 300	9 \pm 6
8	1.6 \pm 0.5	1.6 \pm 1.0	46 \pm 20	7.7 \pm 2.8	NA ⁱ	NA	NA	
16	5.3 \pm 1.2	15 \pm 6.1	7.2 \pm 0.4 ^c	3.9 \pm 0.8	NA	NA	NA	
24	3.4 \pm 2.0	5.3 \pm 0.4 ^c	8.2 \pm 1.0 ^c	5.3 \pm 1.4	NA	NA	NA	
36	3.4 \pm 2.4	3.3 \pm 2.1	2.9 \pm 0.2 ^c	6.4 \pm 2.5	NA	NA	NA	
1 (soluble salt dose) ^e	2000 \pm 610	810 \pm 390	620 \pm 58 ^c	14 \pm 7.9	3400 \pm 2300	1150 \pm 300	530 \pm 400	37 \pm 11
1 (blank)	1.4 \pm 0.2	0.6 \pm 0.4	10 \pm 1.4 ^c	0.8 \pm 0.4	120 \pm 44	33 \pm 28	180 \pm 31	1 \pm 0.1
36 (blank)	2.5 \pm 1.7	28 \pm 0.6 ^c	0.7 \pm 0.2 ^c	2.9 \pm 0.4	NA	NA	NA	

^a Each rabbit dosed with 2.0 g of BMW-1/kg of body weight (average animal weight of 2.1 kg) for an average dose of 4.2 g of BMW-1 (16.4 mg of Pb). ^b Corrected for percent moisture in solids. ^c No error is presented, because only one measurement was obtained. ^d Assuming average rabbit blood volume of 56 mL/kg of body weight (43). ^e Error based on two measurements. ^f No fluid available after centrifugation. ^g One hour between dosing and sample collection. ^h Value calculated by dividing the Pb mass in stomach fluid by total Pb mass in stomach. Associated error calculated as standard deviation of Pb mass in stomach fluid divided by total Pb mass in stomach. ⁱ NA, not analyzed.

not resolved during capillary electrophoresis. Lactic acid, which may be produced in the stomach (37), has been observed previously in the small intestines of rabbits (38). Because the concentrations of acetate, citrate, lactate, and malate were variable throughout the rabbit GI tract, an average concentration of 500 mg/L for each acid was used during the *in vitro* experiments.

Lead in *in Vivo* Fluid and Solid Samples. Lead analyses were performed on all of the fluid samples and on the solid samples collected from the 0.5–6-h post-dose groups. Solid samples from the 8–36-h post-dose groups were not analyzed, because the solid Pb mass had traversed the small intestine and was no longer available for dissolution and absorption (Table III). Reported data were corrected for percent moisture in the solids and are reported as mass (μg) rather than concentration to account for the varying fluid volumes and solid masses present in each rabbit (Table III). Data are presented with standard deviations for each set of three rabbits and indicate substantial interanimal variability (Table III); observed standard deviations are generally in the range of 30–50%. Lead masses in the 1- and 36-h control rabbits are less than or equivalent to any of the dosed rabbits, indicating that background Pb masses are 1–10 and 30–180 μg in GI tract fluids and solids, respectively (Table III), probably due to low concentrations of Pb (0.75 mg/kg) present in the rabbit feed.

Acidification of the small intestinal fluid with HNO_3 , for preservation, resulted in the formation of a flocculent precipitate (believed to be an agglomeration of proteins and enzymes that are present in small intestinal fluid), which was readily resuspended when the pH was raised to >5.0 using NaOH. To evaluate the banding of Pb by the precipitate, three small intestinal fluids were analyzed for Pb concentration, with the precipitate both solubilized and precipitated and removed by centrifugation. Comparison of the Pb concentrations in the fluid samples after solubilization of the precipitate indicates that 63 \pm 4% of the total Pb in solution is removed by this precipitation reaction. Furthermore, a dialysis experiment using 12000 MWCO tubing indicated that 93% of Pb in one small intestinal fluid sample was incapable of diffusing out of the dialysis bag in 24 h. These results suggest that a

substantial portion (approximately 60–95%) of solubilized small intestinal Pb is bound to a protein or enzyme that is >12000 MW.

Mean solubilized Pb masses from BMW-1 as a function of time and GI tract compartment exhibit the expected profile (Table III), with maximum solubilization in the stomach, small intestine, and large intestine occurring at approximately 1.5, 3, and 6 h, respectively. Maximum solubilized Pb masses decrease in each successive segment of the GI tract, with stomach, small intestinal, and large intestinal masses of 0.62, 0.25, and 0.053 mg recovered, respectively. The decrease in Pb mass in solution between stomach and small intestine is probably due to precipitation or sorption reactions that occur as the pH is increased from 1.3 to 7, whereas the decrease in Pb mass between small and large intestine is most likely due to Pb absorption across the intestinal epithelium. Mean Pb masses in GI tract solids are consistent with the pattern observed in the fluid phase (Table III). Emptying of solid Pb from the stomach into the small intestine is complete within 3 h, during which time Pb in the small intestinal solids reaches a maximum (Table III).

When considering available Pb based on either *in vivo* or *in vitro* results, it should be recognized that absorption of Pb in the small intestine is a nonequilibrium process. Four pools of Pb are present in the small intestine: (1) solubilized Pb, either ionic or bound by small organic ligands (Pb_{sol}); (2) solubilized Pb that is bound to large MW entities (Pb_{bnd}); (3) Pb that was solubilized in the stomach but has sorbed to solid surfaces on entering the small intestine (Pb_{orb}); and (4) Pb that was insoluble in the stomach (Pb_{insol}). Because Pb_{insol} is unlikely to be solubilized during passage through the small intestine, this fraction of ingested Pb is not bioaccessible. The other three forms of Pb are all available for absorption. Pb_{sol} is immediately available, while Pb_{bnd} and Pb_{orb} must become unbound or desorbed, respectively, prior to absorption in the small intestine. However, as Pb_{sol} is absorbed, Pb_{bnd} and Pb_{orb} could be released into solution to regain equilibrium partitioning of Pb between the fluid and solid phases. Therefore, assuming that the binding or adsorption mechanisms are reversible, the entire pool of Pb_{sol} , Pb_{bnd} , and Pb_{orb} [which is equal to the total Pb

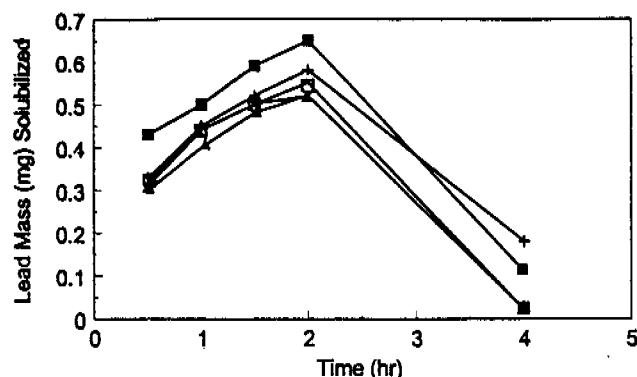


Figure 2. Average Pb mass (mg) solubilized during the *in vitro* method development testing: (■) standard test, (±) rabbit chow present during standard test, (□) no acids present during test, (▲) no enzymes present during test, (×) no enzymes or acids present during test.

in the stomach fluid phase, because surface adsorption of Pb will be negligible at the acidic pH of the stomach (39)] is theoretically available for absorption, if the kinetics of Pb absorption across the intestinal epithelium are sufficiently rapid with respect to intestinal transit time and to the kinetics of Pb release or desorption. The mass of Pb present in the Pb_{bed} and Pb_{orb} pools and the relative release and desorption kinetics were not evaluated during this study. Consequently, the maximum mass of solubilized Pb in the stomach compared to the mass that is insoluble (e.g., the partitioning ratio from solid to fluid) represents a conservative (upper) estimate of available Pb.

Based on the above reasoning, calculation of *in vivo* Pb solubility from BMW-1 (mass of Pb solubilized divided by total Pb present in the stomach) shows that the solubilized Pb fraction increased from an initial 3% to a maximum of 9% at the 1.5-h time point (Table III). The limited Pb dissolution from mine waste is due to the low solubility of Pb-bearing minerals, kinetic dissolution limitations (10), and encapsulation of Pb phases by inert matrices (e.g., silicates, pyrite, and jarosite) (5).

A total of 37% of the recovered Pb from $Pb(OAc)_2$ is present in the stomach fluids after 1 h, suggesting that the balance is present in the stomach solids. It is unclear why the bulk of Pb from $Pb(OAc)_2$ was present in the stomach solids. Blood-lead response 1 h after ingestion of $Pb(OAc)_2$ was 9.4 times larger than from an equivalent mass of Pb contained in BMW-1, supporting maximum *in vivo* mine-waste-lead solubility, relative to $Pb(OAc)_2$, of approximately 10%.

Lead in *in Vitro* Fluids. Reproducibility of the *in vitro* method was tested by conducting triplicate experiments using BMW-1, for which the maximum Pb mass solubilized (2-h sample) was 0.65 ± 0.04 mg. The method development tests, consisting of the addition of rabbit chow, the omission of organic acids, the omission of GI tract enzymes, and the omission of both acids and enzymes (each performed in duplicate), resulted in maximum solubilized Pb masses at the 2-h time point of 0.58 ± 0.04 , 0.55 ± 0.01 , 0.52 ± 0.02 , and 0.52 ± 0.02 mg of Pb, respectively (Figure 2). The blank flasks were below the instrument detection limit (IDL, 0.10 mg/L) in each experiment. The method detection limits (IDL \times dilution factor \times volume in flask) in the mine-waste and method development flasks and the $Pb(OAc)_2$ flask were 0.020 and 0.60 mg, respectively.

The method development experiments indicate that the addition of rabbit chow to the reaction flask causes a slight

decrease in the mass of Pb solubilized in the stomach from 0.65 ± 0.04 to 0.58 ± 0.04 mg, while the mass of Pb in the soluble phase during the small intestinal incubation was increased slightly from 0.11 ± 0.04 to 0.18 ± 0.03 mg (Figure 2). The reason for a decrease in stomach-solubilized Pb in the presence of rabbit chow is unknown, while the increase in intestinal solubilized Pb is most likely due to a solubilized component of rabbit chow, probably citrate, amino acids, or suspended organic matter, that is capable of binding Pb and retaining it in solution. The absence of either organic acids or GI tract enzymes, or both components together, results in a 20% decrease in maximum stomach Pb solubility (0.65 ± 0.04 to 0.52 ± 0.02 mg), consistent with the observation that the presence of organic acids in solution may increase the dissolution of Pb-bearing phases (38). In addition, both acids and enzymes are necessary to retain Pb in solution during the small intestinal phase (Figure 2). These data suggest that organic acids and GI tract enzymes in the pH 7 environment of the small intestine either bind Pb or inhibit the formation of Pb-bearing precipitates.

Based on the reasoning presented in the *in vivo* results section, the fraction of available Pb *in vitro* from BMW-1 ($4 \pm 0.2\%$), determined by dividing the average dissolved Pb mass at 2.0 hr (0.65 mg) by the mass of Pb in the flask (15.6 mg) (Table IV), was in good agreement with the *in vivo* result ($9 \pm 4\%$). The comparison of *in vivo* and *in vitro* solubilized Pb masses (1.5 h for *in vivo* and 2.0 h for *in vitro*) were nearly identical: 0.62 versus 0.65 mg for the *in vivo* and *in vitro* systems, respectively (Tables III and IV). The overall rate of Pb dissolution in the stomach (the rate of change in solubilized Pb with time in Table III versus Table IV) was similar in the *in vivo* and *in vitro* systems, although the *in vitro* dissolution rate was more constant (less variability in the rate of Pb dissolution), indicating that dissolution kinetics are important in controlling Pb bioaccessibility from BMW-1.

The $Pb(OAc)_2$ results indicate that, after correction for Pb emanating from the blank mine waste (100 mg/kg of bulk Pb concentration resulted in 0.07 mg in the *in vitro* test), 68% of the Pb from $Pb(OAc)_2$ is bioaccessible *in vitro* in the presence of a mine-waste matrix, while 76% is available when no matrix is present (Table IV). The discrepancy between available Pb from $Pb(OAc)_2$ during *in vitro* (68 or 76%, depending on the matrix present) and *in vivo* (37%) testing could be due to a variety of factors, including (1) pH differences between the systems, (2) lower concentrations of organic acids *in vivo* than used *in vitro*, or (3) incomplete recovery of ingested material during the *in vivo* study. Soluble Pb mass from $Pb(OAc)_2$ in the absence of mine waste decreased by a factor of 2 in the small intestinal simulation (Table IV), while in the presence of mine waste, soluble Pb decreased by a factor of 9. These results suggest that Pb absorption to the mine-waste surface may reduce Pb solubility in the small intestine.

The mine-waste samples BMW-2-4, consisting predominantly of lead phosphates, manganese-lead oxides, iron-lead oxides, and iron-lead sulfates, also resulted in limited bioaccessible Pb when tested by the *in vitro* method (0.5–6%, Table IV). BMW-2, which consists of ferro-manganese lead oxides and iron-lead sulfate, produced only 0.5% available Pb during the *in vitro* test and reached equilibrium dissolved Pb concentration prior to collection of the initial sample (0.5 h, Table IV). BMW-3 and BMW-4

Table IV. Mass of Pb Solubilized *in Vitro* Compared to *in Vivo* Results

time (h)	Pb mass (mg) solubilized from mine waste during <i>in vitro</i> test					
	BMW-1	BMW-2	BMW-3	BMW-4	Pb(OAc) ₂ /no mine waste	Pb(OAc) ₂ /blank mine waste
0.5	0.43 ± 0.04	0.02 ^a	0.94 ^{a,b}	0.06 ^a	11.0 ± 0.2	10.6 ± 0.2
1.0	0.50 ± 0.02	0.02	1.09 ^b	0.09	11.9 ± 0.2	10.4 ± 0.3
1.5	0.59 ± 0.06	0.02	1.30 ^b	0.11	11.8 ± 0.3	10.2 ± 0.1
2.0	0.65 ± 0.04	0.02	1.26 ^b	0.15	11.4 ± 0.1	10.2 ± 0.1
mass of Pb (mg) solubilized in small intestine simulation	0.11 ± 0.04	<0.008 ^d	0.03	0.04	5.3 ± 0.2	1.0 ± 0.1
mass of Pb added to flask (mg)	15.6	4.1	23.3	7.2	15.6	15.6
fraction available Pb <i>in vitro</i> (%)	4 ^d	0.5 ^d	6 ^d	2 ^d	76 ^d	68 ^d
fraction available Pb <i>in vivo</i> (%)	9 ^e				37 ^e	

^a No standard error available because only one test was conducted. ^b Based on detection limit of 0.1 mg/L of Pb by direct-aspiration atomic absorption spectrometry. ^c Method for establishing maximum available Pb *in vivo* same as *in vitro*, using data from Table II. ^d Fraction of bulk Pb mass solubilized during *in vitro* test (4.0 g substrate/40 mL of fluid). ^e Fraction of bulk Pb mass solubilized during *in vivo* feeding study.

(6% and 2% available Pb, respectively) contain greater percentages of Pb minerals such as lead oxides, lead silicates, and anglesite (Table I), which appear to be more soluble than the lead phosphates, ferromanganese lead oxides, and iron-lead sulfates. In addition, BMW-3 reached equilibrium concentration after only 1.5 h, while BMW-4 did not reach equilibrium during the 2-h stomach incubation. These data indicate that dissolution kinetics limit Pb bioaccessibility from the suite of Pb minerals present in BMW-1 and BMW-4 but not from BMW-2 and BMW-3. No discrete Pb mineral phase appears consistently responsible for the available Pb from these mine wastes, a result that was expected, due to the highly variable Pb mineral composition. Factors such as mineral associations, the presence and type of alteration and precipitation rinds, encapsulation of Pb phases in inert minerals (e.g., silicates), and the presence of surface-adsorbed Pb are likely to be of equal or greater importance in controlling Pb bioaccessibility in the GI tract.

Conclusions

This study has demonstrated that an *in vitro* method can be an effective tool in evaluating the geochemical and physiological factors controlling the dissolution of mine-waste Pb in the GI tract.

The rabbit was determined to be an appropriate model for human GI tract conditions, with respect to pH and transit times. In a validation study using female New Zealand White rabbits, average pH values were 1.3 and 3–4 in the stomach under fasting and fed conditions, respectively, and approximately 7 in the small intestine, similar to pH values reported in humans. Acetate, citrate, lactate, and malate are all present in varying concentrations in both human and rabbit GI tracts and may chelate Pb with relative complex formation strengths of citrate > malate > acetate > lactate, thereby increasing Pb solubility and subsequent absorption across the intestinal epithelium by passive diffusion. However, a variety of potential chelating agents (organic acids, amino acids, proteins, enzymes, and tannins) will compete for soluble Pb, and on the basis of two preliminary tests, 60–95% of solubilized Pb in the small intestine appears to be bound to a protein or enzyme >12000 MW. Although a stomach pH of 1.3 was used in the *in vitro* test to maximize Pb dissolution, this stomach acidity would apply only to fasting children. A stomach pH of 3–4, typical when food is in the stomach, would be expected to further reduce Pb bioaccessibility by decreasing the extent of Pb mineral dissolution.

The *in vitro* beaker assay, modeled after an assay method used to determine available Fe from food, was successful in replicating Pb dissolution during the feeding study. Four mine-waste samples with highly variable mineral composition (BMW-1–4) resulted in only 4, 0.5, 6, and 2% bioaccessible Pb, respectively, suggesting that geochemical and physiological factors limit the solubility and, therefore, the availability of Pb from mine waste, regardless of whether the bulk Pb mineralogy consists of the sulfate/sulfide or the oxide/phosphate assemblage. The 4% available Pb from BMW-1 is in good agreement with a 30-day feeding study in Sprague-Dawley rats, using this mine waste, which produced only 3% absolute Pb bioavailability based on blood-lead data (3). In addition, method development testing indicates that although HCl concentration is the most important GI component controlling Pb dissolution in the stomach, both organic acids and enzymes are necessary to retain Pb in solution during the small intestinal incubation. Based on these results, the assay appears to provide a useful, rapid, screening-level test to predict maximum available Pb from mine wastes.

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